

Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
L1	1748	544/235, 514/248	US-PGPUB; USPAT	OR	OFF	2005/10/07 13:34

KM
Day : Friday
Date: 10/7/2005
Time: 13:06:42



PALM INTRANET

Inventor Information for 10/799404

Inventor Name	City	State/Country
FU, JIAN-MIN	BURNABY	CANADA

[Appln Info](#) [Contents](#) [Petition Info](#) [Atty/Agent Info](#) [Continuity Data](#) [Foreign Data](#)

Search Another: Application# or Patent#
PCT / / or PG PUBS #
Attorney Docket #
Bar Code #

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Back to [PALM | ASSIGNMENT | OASIS | Home page](#)

10/799,404

Page 3

Broad search
for 10/799,404
10/799,406
10/799,407

chain nodes :

17 18 20 21 22

ring nodes :

1 2 3 4 5 6 7 8 9 10 11 12 13 14 15

chain bonds :

4-22 9-10 11-20

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-9 7-8 8-9 10-11 10-15 11-12 12-13 13-14
14-15

exact/norm bonds :

1-2 1-6 2-3 3-4 4-5 4-22 5-6 5-7 6-9 7-8 8-9 9-10 10-11 10-15 11-12
11-20 12-13 13-14 14-15

isolated ring systems :

containing 1 : 10 :

G1:C,N

G2:CH3,X

Match level :

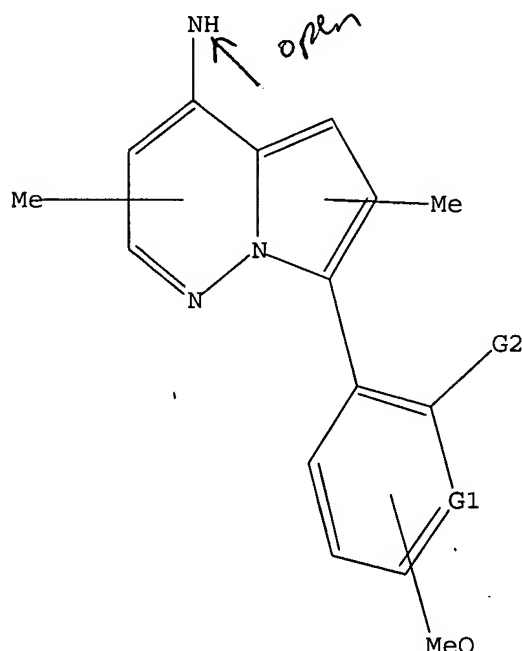
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom
11:Atom 12:Atom 13:Atom 14:Atom 15:Atom 17:CLASS 18:CLASS 20:CLASS 21:CLASS
22:CLASS 23:CLASS 24:CLASS 25:CLASS

L1 STRUCTURE UPLOADED

=> d 11

L1 HAS NO ANSWERS

L1 STR



G1 C,N

G2 Me,X

<10/07/2005>

Habte

Structure attributes must be viewed using STN Express query preparation.

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=> s 11
SAMPLE SEARCH INITIATED 09:24:19 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED -          7 TO ITERATE

100.0% PROCESSED      7 ITERATIONS          0 ANSWERS
SEARCH TIME: 00.00.01
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FULL FILE PROJECTIONS:	ONLINE	**COMPLETE**
	BATCH	**COMPLETE**
PROJECTED ITERATIONS:	7 TO	298
PROJECTED ANSWERS:	0 TO	0

L2 0 SEA SSS SAM L1

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=> s 11 sss full
FULL SEARCH INITIATED 09:24:28 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED -          87 TO ITERATE
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100.0% PROCESSED	87 ITERATIONS	8 ANSWERS
SEARCH TIME:	00.00.01	

L3 8 SEA SSS FUL L1

=> file caplus		SINCE FILE	TOTAL
COST IN U.S. DOLLARS		ENTRY	SESSION
FULL ESTIMATED COST		161.33	161.54

FILE 'CAPLUS' ENTERED AT 09:24:33 ON 07 OCT 2005
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FILE COVERS 1907 - 7 Oct 2005 VOL 143 ISS 16
 FILE LAST UPDATED: 6 Oct 2005 (20051006/ED)

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This file contains CAS Registry Numbers for easy and accurate substance identification.

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L4 3 L3

<10/07/2005> Habte

10/799,404

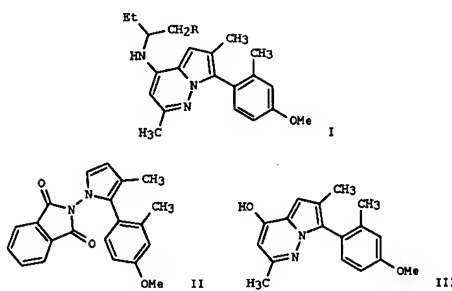
Page 5

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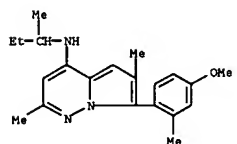
L4 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2005 ACS ON STN
ACCESSION NUMBER: 2004:857604 CAPLUS
DOCUMENT NUMBER: 141:332205
TITLE: Preparation of pyrrole[1,2-b]pyridazine compounds as
CRF receptor antagonists for the treatment of
disorders such as anxiety and depression
INVENTOR(S): Fu, Jian-min
PATENT ASSIGNEE(S): Pharmacia & Upjohn Company, USA
SOURCE: PCT Int. Appl., 26 pp.
CODEN: PIIX02
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2004087708	A1	20041014	WO 2004-1B1006	20040322
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KK, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NA, NL, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, US, UZ, VC, VN, YU, ZA, ZM, ZR, BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AH, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, DE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
US 2004209887	A1	20041021	US 2004-792940A	20040312
PRIORITY APPLN. INFO.: US 2003-460698P P 20030404				
C1				



L4 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

RN 773086-73-0 **CAPLUS**
CN Pyrrol[1,2-b]pyridazin-4-amine, 7-(4-methoxy-2-methylphenyl)-2,6-dimethyl-1-methyl-4-((dimethylamino)methyl)-; N-((1-methylpropyl)- (SCI) (CA INDEX NAME)



REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE REFORMAT

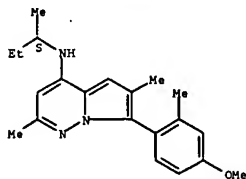
L4 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

AB Disclosed are novel CRF receptor antagonists and their use in the treatment of a variety of disorders, including disorders manifesting hypersecretion of CRF, or associated with CRF or CRF receptors, such as anxiety, and depression. The CRF receptor antagonists of the invention have the structure of formula I (R = H or Me), including stereoisomers or mixts. of stereoisomers, pharmaceutically acceptable salts, and pharmaceutically acceptable esters. Compds. I were tested in several bioassays, and had IC50 values of less than 3 nM in a CRF1 receptor binding assay. For example, 4-bromo-3-methylanisole was treated with t-BuLi followed by reaction with α -methyl- γ -butyrolactone to give a ring-opened hydroxy ketone which underwent Swern oxidation to yield the corresponding formyl ketone. This dicarbonyl compound was cyclized with N-aminophthalimide to afford pyrrole II, which was deprotection with hydrazine and then converted to hydroxycyclohexene III via cyclocondensation with Et trans-3-thioxocrotonate. Bromination of III with PBz3 followed by amination of the resulting bromide with (S)-sec-butylamine led to pyrrole[1,2-b]pyridazine (S)-I (R = H). Claimed uses also include (1) use of labeled compds. I in competitive binding assays for screening of other CRF receptor ligands, and (2) use of labeled I for detecting CRF receptors in tissue.

IT 773086-71-EP 773086-72-9P 773086-73-0P
RL: ARG (Analytical reagent use); BSU (Biological study, unclassified);
PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); ANST (Analytical study); BIOL (Biological study); PREP
(Preparation); USES (Uses)
(drug candidate; preparation of pyrrolopyridazine derivs. as CRF receptor
antagonists)

RN 773086-71-8 CAPLUS
CN Pyrrole[1,2-b]pyridazin-4-amine, 7-(4-methoxy-2-methylphenyl)-2,6-dimethyl-N-[(1S)-1-methylpropyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



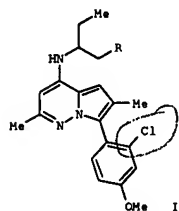
RN 773086-72-9 CAPLUS
CN Pyrrolol[1,2-b]pyridazin-4-amine, N-(1-ethylpropyl)-7-(4-methoxy-2-methylphenyl)-2,6-dimethyl- (9CI) (CA INDEX NAME)

L4 ANSWER 2 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER: A2004-057173 CAPLUS

ACCESSION NUMBER: 2004:851713 CAPLUS
DOCUMENT NUMBER: 141:350182
TITLE: Preparation of pyrrolo[1,2-b]pyridazine compounds and their use as GPR55 receptor antagonists

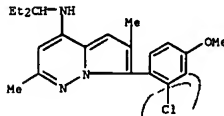
INVENTOR(S): Fu, Jian-min
PATENT ASSIGNEE(S): Fuzhou Inc., USA
SOURCE: U.S. Pat. Appl. Publ., 12 pp.
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2004204415	A1	20041014	US 2004-799407	20040312
WO 200407709	A1	20041014	PCT/US-2004-18951	20040322
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, FR, GE, GH, GR, HI, HK, IL, IN, IS, JT, KE, KG, KP, KR, LC, LK, LS, LT, LV, MA, MD, MG, MK, MN, MW, NA, NI, NO, NZ, OM, PH, PL, PT, RO, RU, SG, SD, SE, SG, SK, SL, SY, TZ, TM, TN, TR, TZ, UA, US, UZ, VC, VN, YU, ZA, ZH, RW: BW, GH, GM, KE, LS, MW, MZ, SD, SI, SZ, TZ, UG, ZW, AH, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CH, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
SECOND APPN. INFO.: HER SOURCE(S):		US 2003-460734P	P 20030404	
MARPAT 141:350182				



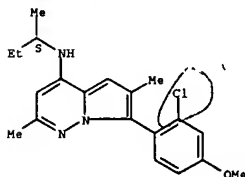
AB The title compds. [I : $R = H, Me$], useful in the treatment of a variety of disorders, including disorders manifesting hypersecretion of CRF or associated with CRF or CRF receptors, such as anxiety, and depression, were prepared. E.g., a multi-step synthesis of I [$R = Me$], starting from 4-bromo-3-chloroanisole and α -methyl- β -hydroxybutyrolactone, was given. The compds. I showed K_1 of <2.0 nM in *in vitro* CRF receptor given. The compds. I showed K_1 of <2.0 nM in *in vitro* CRF receptor

- L4 ANSWER 2 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)
binding assay. The pharmaceutical compn. comprising the compd. I is claimed.
- IT 775345-59-0 775345-60-3P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
(preparation of pyrrolo[1,2-b]pyridazine compds. and their use as CRF receptor antagonists)
- RN 775345-59-0 CAPLUS
- CN Pyrrolo[1,2-b]pyridazine-4-amine, 7-(2-chloro-4-methoxyphenyl)-N-(1-ethylpropyl)-2,6-dimethyl- (9CI) (CA INDEX NAME)

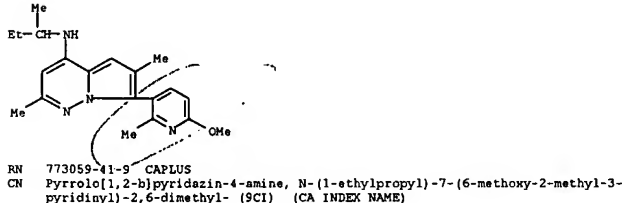


RN 775345-60-3 CAPLUS
CN Pyrrolo[1,2-b]pyridazine-4-amine, 7-(2-chloro-4-methoxyphenyl)-2,6-dimethyl-N-(1S)-1-methylpropyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

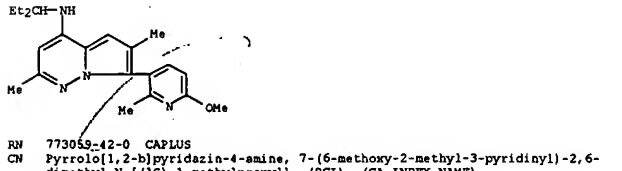


- L4 ANSWER 3 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)
hypersecretion of CRF or assoccd. with CRF or CRF receptors, e.g. anxiety and depression. CRF receptor antagonists of the invention have structure I (R = H, Me), including stereoisomers or mixts. of stereoisomers, pharmaceutically acceptable prodrugs, or pharmaceutically acceptable salts thereof.
- IT 773059-40-8 773059-41-9 773059-42-0
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(pyrrolopyridazine compound CRF receptor antagonists, and use in treatment of CRF- and CRF receptor-associated disorders)
- RN 773059-40-8 CAPLUS
- CN Pyrrolo[1,2-b]pyridazine-4-amine, 7-(6-methoxy-2-methyl-3-pyridinyl)-2,6-dimethyl-N-(1-methylpropyl)- (9CI) (CA INDEX NAME)



RN 773059-41-9 CAPLUS

CN Pyrrolo[1,2-b]pyridazine-4-amine, N-(1-ethylpropyl)-7-(6-methoxy-2-methyl-3-pyridinyl)-2,6-dimethyl- (9CI) (CA INDEX NAME)



RN 773059-42-0 CAPLUS
CN Pyrrolo[1,2-b]pyridazine-4-amine, 7-(6-methoxy-2-methyl-3-pyridinyl)-2,6-dimethyl-N-(1S)-1-methylpropyl- (9CI) (CA INDEX NAME)

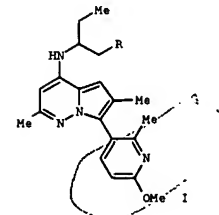
Absolute stereochemistry.

- L4 ANSWER 3 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)
ACCESSION NUMBER: 2004:857172 CAPLUS
DOCUMENT NUMBER: 141:325761
TITLE: Pyrrolo[1,2-b]pyridazine compound corticotropin-releasing factor (CRF) receptor antagonists and their use in the treatment of CRF- and CRF receptor-associated disorders
- INVENTOR(S): Fu, Jian-min
PATENT ASSIGNEE(S): Pfizer Inc., USA
SOURCE: U.S. Pat. Appl. Publ., 11 pp.
CODEN: USXOXC0
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2004204414	A1	20041014	US 2004-799406	20040312
WO 2004087710	A1	20041014	WO 2004-18971	20040322

V: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
RW: BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG

PRIORITY APPLN. INFO.: US 2003-459744P P 20030402
GI



AB The invention discloses CRF receptor antagonists and their use as treatment of a variety of disorders, including disorders manifesting

- L4 ANSWER 3 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

